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Description of a central Amazonian and Guianan tree frog, genus Osteocephalus (Anura, Hylidae), with oophagous tadpoles

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A new species of the hylid frog genus Osteocepholus is described from Manus, Amazonas, Brazil II is a medium-sized species that resembles O, tourins, but is smaller and has shorter hind legs. In contrast to other species of Osteocepholus, the dornal slarit is barely securitly during the other species of Osteocepholus, the dornal slarit is barely securitly during the other species of Osteocepholus, the dornal slarit is barely securitly during the other species of Osteocepholus, the dornal slarit is barely security during a balant of the other species of the other species

Most species of the neotropical hylid frog genus Osteocephalus Steindachner, 1862 are identifiable by the presence of well-ossified skulls, the presence of paired lateral vocal sacs, and rugose or warty dorsal skin in males and relatively smooth skin in females. TAUB & DUELLMAN (1971) reviewed and defined the genus and recognized five species: O. buckley (Boulenger, 1882), O. leprieuri (Dumčri & Bibron, 1841), O. pearsoni (Gaige, 1929), O. taurinus Steindachner, 1862 and O. verruciger (Werner, 1901) (verrucigerus auct.). DUELLMAN (1974) followed Cope (1867) in placing Hyla langsdorffii Duméril & Bibron, 1841 in the genus. Topotypes of Hyla elkejungingerae Henle, 1981 were identified as O. verruciger by DUELMAN (in litt. to W. BÖHME, pers. comm.) but placed into the synonymy of O. taurinus by HOGOMODE (1982) MARTINS & CAROSO (1987) described O. subrilis from Acre, Brazil, and DUELLMAN & HOGOMODE (1992) placed Hyla rodriguezi Rivero, 1968 in Osteocephalus. AVARZAGUENA et al. (1992a) described five species, O. aecli, O. edelcar, O. galani, O. laucelabris and O. rimarum from Venezuelan table mountains and placed them in the O. rodriguezi group. AVARZAGUENA et al. (1992b) transferred this group to a



new genus, Tepuihyla. Thus, between 7 and 14 described species are currently comprised in the genus.

An additional species from midwestern Amazonia is in the process of being described by M. MARTINS and M. GORDO (pers. comm.). One more species from central Amazonia and two Guyanese states has been known for much more than a decade and has appeared in the literature many times as Osteocephalus sp. (HOCMORD, 1979; ZIMMERMAN, 1983; ZIMMERMAN & BOCART, 1984; MARTINS & CARDOSO, 1987; FIERO, 1990; HODL, 1990, 1993; ZIMMERMAN & BOCART, 1984; MARTINS & CARDOSO, 1987; FIERO, 1990; HODL, 1990, 1993; ZIMMERMAN & BOCART, 1980; HOCGMOED & AVIL-PIRES, 1991; DUELLMAN & HOCGMOED, 1992; WEYCOLDT & JUNGFER, 1993) or erroneously as Osteocephalus buckleyi (ZIMMERMAN & BOCART, 1988). In the course of our independent work on the reproductive biology and tadpole morphology of this species, we felt that the taxonomic status of the frog needed to be resolved. Therefore we describe it here as

Osteocephalus oophagus sp. nov.

Holotype. – MZUSP 69852, an adult male, collected by K.-H. JUNGFER on 9 April 1993 at Reserva Florestal Adolfo Ducke (2°55′S, 59°59′W), situated at km 26 of the Rodovia AM-010 (Manaus-Itacoatiara), Estado do Amazonas, Brazil.

Paratypes. – NMW 32925.1-2, collected by W. Hön. on 28 February 1978; MPEG 4845-4846, collected by M. J. HENZL, L. S. FORD and A. LIMA on 9 February 1992; AMNH A.13618-136184, collected by M. J. HENZL and L. S. FORD on 3 March 1992; MZUSP 69853, INPA 01446 and 01448, SMNS 10801-10802, ZFMK 57137-57138, all collected by K.-H. JUNGER between January and April 1993; all from Reserva Florestal Adolfo Ducke. For sexes and Museum abbreviations see Table I.

Definition. - For purposes of comparisons, we follow TBUEB & DUELLMAN's (1971) standards. A medium sized species of Osteocephalus (maximum known snout-vent length (SVL) 47.2 mm in males, 62.7 mm in females). Dorsal skin in males with a few non-spinous tubercles or smooth, in females smooth. Skin on flanks smooth, slightly shagreened posterior to the insertion of the arm in some specimens. Webbing on hand moderate, up to first third of antepenultimate phalans on inner edge of third finger. Dorsum brown to grey with or without irregular tan fleeks, spots, mottling or reticulation, with white spots in some specimens. Venter white; in some specimens very light brown reticulation present on the chest. Lips brown or grey, with or without a cream subocular spot extending to the edge of the tympanum. Flanks brown or grey. White mottling or reticulation no posterior half of the flanks; many white spots on dark ground present in some specimens.

Diagnosis. — Ozteocephalus oophagus is distinguished from the frogs of the O. rodriguezi group by substantial webbing on the hand (absent or rudimentary in the O. rodriguezi group), and from O. huckleyi by the lack of a conspicuous row of tubercles on the tarsus and prominent supraorbital tubercles. O. verruciger and O. elkejungingerae are larger and the dorsum of males is covered by many spinous tubercles. O. leprievati has less webbing on the hands (web reaching base of antepenultimate phalanx on inner side of third finger) and in some cases transverse lines or bars on the dorsum that are absent in O. oophagus. feet that are lacking in O. oophagus. The new species is distinguished from O. pearsoni by the lack of brown reticulation on the venter (present only in chest area, if at all, in O. oophagus) and more webbing on the hands (to base of antepenultimate phalanx on inner side of third finger in O. pearsoni). O. subtilis has a black iris (golden with black rays in O. oophagus) and a glandular line above the vent (absent in O. oophagus). An additional undescribed species from midwestern Amazonia bears a conspicuous, complete white line running around the whole of the upper lip (faint and interrupted in O. oophagus), and its supratympanic fold is smooth (tubercular in O. oophagus) (M. MARTINS, pers. comm.). O. oophagus is most easily confused with O. taurinus, with which it shares the golden iris with radiating black lines. O. taurinus is a larger frog that reaches 104 mm. The dorsum in males is covered by many spinous tubercles with keratinized tips. The webbing on the hands is slightly more extensive than in O. oophagus, reaching the middle of the antepenultimate phalanx on the inner side of the third finger. The legs are longer than in O. oophagus. When stretched forward and angled at 90° at the tibiotarsal articulation, the tarsus reaches the tip of the snout in O. oophagus and extends beyond that point in O. taurinus. Furthermore, adults of O, taurinus possess two elevated longitudinal bony ridges (frontoparietal flanges) in the interorbital area, that are not visible in O. oophagus, but may be felt in large specimens when rubbing with the fingertips.

Description of holotype. - Measurements and proportions (following DUELLMAN, 1970) are given in Tables I and II. A male of 43.8 mm SVL (fig. 1a). Body wider than the head. Diameter of the tympanum slightly wider than half the eve diameter. Head flat between orbits, slightly concave in the intercanthal region and truncate in lateral and dorsal aspect. Nostrils elevated, internarial region slightly depressed. Canthus rostralis slightly rounded, loreal region concave, bearing a few rounded warts below the canthus. An elevated dermal fold ascending posterior to the mid level of the eve to the area above the tympanum and sloping to the insertion of the arm. Tympanum distinct and rounded. A weak axillary membrane extending to less than one fifth the length of the humerus. A row of flat, barely raised warts visible on the ventrolateral edge of the forearm, extending onto the proximal half of the fourth finger. Fingers with moderately large discs, that of the third finger about four fifths the diameter of the tympanum. Enlarged prepollex bearing an elliptical tubercle. Nuptial pads absent (dark brown before), as the frog was no more in breeding condition when preserved on 28 July 1993. Distal subpalmar tubercle bifid on finger IV. Webbing basal between finger I and II and with a webbing formula (SAVAGE & HEYER, 1967; MYERS & DUELLMAN, 1982) of II1.5-3⁻III2.5-2⁺IV in the others. Relative finger lengths of adpressed fingers 3>4>2>1 (fig. 2a). Legs relatively short. Tarsus reaching the tip of the snout when stretched forward parallel to the body axis and bent at 90° at the tibiotarsal articulation. Inner metatarsal tubercle large, flat and elliptical. No outer metatarsal tubercle. Toe webbing formula 11⁺-2⁻III⁺-2IIII⁺-2IV2-1⁺V. Relative toe lengths 4>5>3>2>1 (fig. 2b). Vent opening positioned posteriorly at the upper level of the thighs.

Dorsally, skin weakly granular with low tubercles in the inter- and postorbital region and few on the anterior part of the dorsum. Dorsal aspects of the arms weakly granular and those of the legs smooth. Laterally, skin tubercular posteroventral to the tympanum, shagreened on the anterior half of the flanks and smooth on the posterior half, tubercular

Table I. - Measurements of type specimens of Osteocephalus oophagus. Measurements in mm: ED, eye diameter; FL, foot length; HL, head length; HW, head width; SVL, snout-vent length; TL, tiba length; TD, tympanum diameter. Museum abbreviations: AMNH, a da Amazônia, Manaus; MPEG, Muasu Paraense Emilio Goeldi, Beldm; M2USP, Muasu de Zoologia, Universidade e São Paulo: NNWY, Naturhistorisches Museum Wier; SMNS, Staatliches Museum für Naturkunde, Stutgart; ZFMK, Zoologisches Forschungsinstitut und Museum Alexander Koenig, Bonn.

	SVL	HL	нw	ED	TD	TL	FL
Males							
SMNS 10802	35.7	12.2	10.7	3.90	2.16	19.5	13.8
MPEG 4846	41.6	14.5	13.7	4.80	3.12	23.5	17.5
INPA 1448	42.3	14.2	13.1	5.10	2.82	22.8	15.8
AMNH 136184	42.5	15.2	13.8	5.10	2.86	24.2	17.8
NMW 32925.2	43.2	14.6	13.9	4.86	2.82	22.7	17.0
ZFMK 57138	43.3	15.5	14.5	5.22	3.24	24.3	17.9
MZUSP 69852	43.8	14.7	13.0	4.56	2.52	23.3	17.2
SMNS 10801	47.2	16.6	14.6	6.00	3.66	23.5	18.3
Females							
AMNH 136183	46.2	16.0	13.7	5.34	3.20	23.8	18,1
ZFMK 57137	49.8	17.2	16.9	5.34	3.48	27.7	20.3
MPEG 4845	53.2	18.2	16.9	5.52	3.96	29.5	22.5
MZUSP 69853	53.6	17.9	16.7	5.82	3.90	30.8	21.2
NMW 32925.1	54.6	18.9	16.5	5,46	4.32	31.2	22.0
INPA 1446	55.6	17.7	16.8	5.10	3.72	29.5	21.6

Table II. - Proportions of male and female Osteocephalus oophagus. Abbreviations: see Table I.

Locality, sex	n		HL/SVL	HW / SVL	TD / ED	TL / SVL	FL / SVL
Reserva Ducke, males	8						
		min.	0.336	0.297	0.553	0.498	0.374
		max.	0.358	0.335	0.650	0.569	0.421
		mean	0.346	0.316	0.585	0.542	0.399
Reserva Ducke, females	6						
		min.	0.318	0.297	0.599	0.515	0.388
		max.	0.346	0.339	0.791	0.575	0.423
		mean	0.339	0.312	0.693	0.551	0.402
Rio Urucu, females	2						
		min.	0.348	0.297	0.735	0.557	0.403
		max.	0.357	0.301	0.835	0.574	0.418
		mean	0.353	0.299	0.785	0.566	0.411

below the vent. Skin on the belly and the posteroventral thigh surfaces granular, the other ventral surfaces smooth. Tongue round. Prevomers angular, with 8 and 10 prevomerine teeth. Vocal slits extending postero-laterally from the middle of the tongue. Vocal sac subgular, median and weakly distensible during call.

Colour in alcohol. – Dorsal surfaces brown with tan flecks and blotches. Flanks light brown with brown motiling on the posterior half. Upper and lower lips bordered by faint creamy stripes often interrupted by tan spots. Venter creamy with some faint light brown motiling on the chest. Throat creamy while with light brown mottling. Ventral surfaces of the arms creamy with a brown hue, those of the legs pale light brown. Posterior thigh surfaces brown. Bones green and visible through the skin. Iris yellowish white with many radiant black stripes and a horizontal black bar on each side of the pupil.

Colour in life. - Depending on illumination and colour of the substrate the frog was sitting on, the dorsal colour varied from light brown with indistinct darker flecks and blotches to dark brown with almost black flecks and blotches. Flanks were brown to creamy brown with darker brown-mottling on the posterior half. Ventral surfaces were white. Bones were green and the iris golden yellow with radiant black stripes and horizontal black bars.

Variation. — The largest male of the type series has an SVL of 47.2 mm, the largest female of 55.6 mm. Thirty marked and released males from the type locality bearing nuptial pade ranged from 35.9 to 45.5 mm, and seventeen ovigerous females from 49.8 to 60.6 mm. One female (INPA 01442) from the Rio Urucu is 62.7 mm. There is little variation in proportions between the sexes, except that the tympana of females are slightly larger than those of males (Table II). The dorsal skin varies from smooth to weakly granular, bearing none or a few more tubercles on the tandency of males to bear a few more tubercles on head and dorsum than females, but some males have both surfaces smooth, while some females have a few tubercles on the head and sometimes on the dorsum. This variation is found in the preserved material at hand, and was also obvious in the many live frogs we saw. Dorsal tuberculation is not a reliable sexual dimorphic character. Breeding males bear horny dark brown nuptial pads on the prepolicies.

A lot of variation was observed in coloration. In preservative, the dorsal colour ranges from grey to dark brown with or without darker blotches or flecks (figs. la-b). Also, the amount of cream spots or motiling dorsally, laterally and around the vent is variable (figs. lc-d). The ventral brown motiling on chest and throat may be lacking or reduced to the area under the lower jaw.

Habitat. - Reserva Florestal Adolfo Ducke, a reserve managed by the Instituto Nacional de Pesquisas da Amazônia (INPA), consists of hilly terra firme lowland forest at an altitude of about 50 m. Frogs usually migrate at night on vertical stems of young trees and bushes normally less than 2 cm in diameter at heights of 0.5 to 2 m. Males call from near or inside phytotelmes.

Reproductive biology. — Pairs of O. oophagus deposit their eggs in phytotelmes, such as epiphytic or ground bromeliads, Buriti Palm (Mauritia flexuosa) leaf axils, water-filled palm bracts lying on the ground or tree holes up to heights of about 35 m. Females return to the deposition sites regularly in intervals of about five days, usually clasped by the same

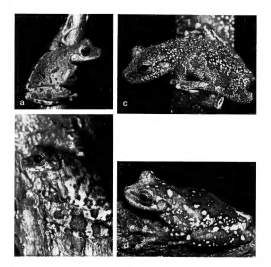


Fig. 1. - Osteocephalus oophagus sp. nov. Specimens from the type locality. (a) Holotype MZUSP 69852, adult male. (b) Male showing numerous tan dorsal blotches. (c) Subadult female with numerous small white spots. (d) Adult female showing extreme extent of lateral and dorsal white spots.

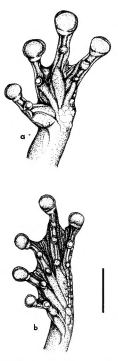


Fig. 2. - Hand (a) and foot (b) of a male Osteocephalus oophagus (paratype INPA 01448). Line equals 5 mm.

males. If there are larvae already present, they consume the newly laid eggs. The eggs not eaten hatch. The larvae starve if the mother fails to return and if they are not provided with trophic eggs (WEYGOLDT & JUNGFER, 1993). A detailed study on the reproductive biology is in progress.

Calls. – The advertisement call usually consists of one to six croaking notes emitted at night and occasionally during the day (type A). At night it is very often followed by one to four (usually two) distinctly different notes that sound like "ka kå" (type B).

Type A of frogs from the INPA-WWF MCSEP reserves and from Parque Nacional Tapajös were described in detail by ZIMMERMAN & BOGART (1984), who recorded an emphasized frequency of 1.62 ± 0.65 kHz, low frequency range of 0.89 ± 0.33 kHz, high frequency range of 2.84 ± 0.42 kHz, and call duration per note of 0.24 ± 0.07 s. ZIMMERMAN & BOGART (1988) again described calls and calling of this frog, under the name of *Osteocephalus buckleyi*. They noted the high intraspecific variability and other characteristics of the call.

Description of the tadpole. - Tadpoles were collected by L. C. SCHESARI on 5 March 1993 in water-filled plastic basins used as egg-laying sites by O. oophague near the Igarapé Acará, Reserva Florestal Adolfo Ducke, and preserved in 10 % formalin. Two tadpoles were raised until metamorphosis. Measurements were taken with the optical measuring unit Wild MMS 235. Drawings (figs. 3-4) were made with a camera lucida attached to a stereomicroscope. Measurements were made according to GRILITSCH et al. (1993). Developmental stages were determined following GOSNER (1960). Labial tooth row formula (LTRF) is after ALTIC (1970). The following description is based on 10 tadpoles ranging from stage 35 to stage 38. Measurements of larvae are given in Table III.

Tadpole of ORTON's (1953) type IV. Body slightly depressed, ovoid in dorsal view. Snout nearly truncate from above and, in profile, acutely rounded. Eyes positioned dorsolaterally; interorbital distance 1.5-1.7 times the internarial distance, which in turn equals width of oral disc. Nares rimmed, rounded, and directed anterolaterally. Their distance to the eyes about half of their distance to tip of the snout. Spiracular tube sinistral, ventrolateral, and directed posterodorsally. Spiracular opening slightly posterior to mid length of body, to which it is tightly attached. Dorsal and ventral margin of caudal musculature parallel in proximal third, then gradually narrowing and almost reaching tip of the tail. Ventral fin slightly lower than dorsal one, fairly paralleling ventral margin of caudal musculature. Dorsal fin extending a short distance onto body. Oral disc anteroventral and not emarginate. One row of moderately sized marginal papillae with a medial gap in upper labium which may be visible in dorsal view. Submarginal papillae in some specimens in one discontinuous row in lower labium as well as scattered in ventrolateral portion of oral disc. Two rows of denticles on the upper labium, the second one showing a distinct median gap, and three rows on the lower one (LTRF 2(2)/3). Upper jaw sheath arched, lower one V-shaped, both black and without serrations (magnification 50 ×). Colour of measured tadpoles chestnut brown dorsally; venter and caudal musculature lighter.

Ontogenetic change. - Newly metamorphosed frogs 12-13 mm in SVL and completely different in colour from the adults. Dorsal and lateral surfaces grey except for a black canthal stripe continuing as a supratympanic stripe posterior to eye to insertion of the arm. Dorsal surfaces of upper arm and proximal half of lower arm white. An orange spot on

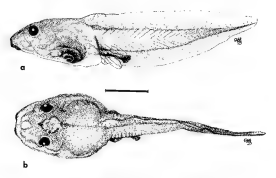


Fig 3 - Dorsal (a) and lateral (b) view of a tadpole of Osteocephalus oophagus in stage 36 (GOSNER, 1960) of typical proportions Line equals 5 mm.

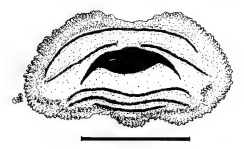


Fig. 4. - Oral disc of a tadpole of Osteocephalus oophagus (same individual as in fig. 3). Line equals 1.4 mm.

Table III. - Measurements of larvae of Osteocephalus oophagus (mean values, ranges in parentheses) Measurements in mm: DT, maximum height of tail: LF, maximum height of lower (ventral) tail fm; SVL, snou-vent length; TL, total length; UF, maximum height of upper (docsa) lail fm; YT, tail length.

Stage	n	TL	SVL	VT	DT	UF	LF
19	t	4 66					
20	4	4.73 (4.35-5 03)					
23	5	5 98 (5 70-6 48)					
27	1	167	7.2	9.5	3 71	1.20	1,19
28	1	19.5	8.6	10.9	4 68	1.61	1 50
30	1	213	9.1	12.2	4.97	1.60	1 62
31	1	24 2	10 2	14.0	5.21	1 61	1 58
33	2	26.2 (25.4-27 0)	117 (116-11.8)	14 5 (13 8-15 2)	6.02 (5.93-6.10)	1 98 (1.86-2.10)	1.76 (1 76-1.76)
34	2	23 7 (22.5-24.9)	10.8 (10.1-11.4)	12 9(12.4-13 5)	5 83 (5.31-6 34)	1 74(1.59-1.89)	1.81 (1.73-1.88)
35	5	27 0 (25 0-28 2)	12.5 (10 0-13.3)	14.5 (12.8 16.0)	6.14 (5.84-6.44)	1 99 (1 90-2 14)	1.84 (1.66-2 02)
36	1	28 9	14 2	14 7	6 49	2.08	175
37	1	27 8	12.9	14.9	6.55	2.08	1.98
38	3	29.5 (26.4 31.6)	13.9 (13 6-14 5)	15.5 (11 9-17 8)	6 44 (6 44-6 44)	2.06 (1.98-2.13)	1.94 (1 88-1 99)
39	3	30 0 (28.4-31.4)	13.8 (13 4-14 3)	16.2 (15.0-17 6)	6.96 (6.27-7.85)	2 07 (1.84-2.35)	1.91 (1.60-2 16)
40	2	34.6 (32.9-36.2)	15.1 (15.0-15.1)	19.5 (17.8-21 2)	6.78 (6.65-6.90)	1.96 (1.57-2.35)	1.99 (1 95-2 04)

elbow, distal dorsal surface of lower arm and proximal dorsal surface of fourth finger. A large white spot, capped by a smaller orange one, on heel. Another orange spot on knee and in some specimens on outer edge of the metatarsus. Finger- and toe-discs orange. Iris bright red without radiating black stripes With this colour pattern, they are virtually indistinguishable from sympatric O. taurinus (juveniles described as O. taurinus by DUELLMAN & LESCURE, 1973, are most likely those of Hyla geographica).

A specimen of 21 mm SVL already had the typical adult pattern, except that the white spot on the upper arm and proximal lower arm was still present. The finger- and toe-dises, as well as the ins, already with black rays and horizontal bars, had an orange hue.

Distribution. — Apart from the Reserva Ducke, about 25 km north of Manaus, we have heard the distinctive call of this species in the forest of the Universidade do Amazonas campus within the city of Manaus and in Reserva Florestal Walter Egler, about 50 km north-northeast of Manaus. The frog is well known from the INPA-WWF Minimal Critical Size of Ecosystems Project (MCSEP) reserves, approximately 80 km northeast of Manaus (ZMMERMAN, 1983; ZMMERMAN & BOGART, 1984, 1988, ZMMERMAN & RODRI-GUES, 1990). HOOGMOED (1979) listed it from "castern Guiana" and HOOGMOED & AVILA-PRES (1991) recorded it from Pett Saut, French Guyana, and remarked that it was also known from Suriname and Brazi. Those records are all in the Guianan biogeographical region (see HOOGMOED, 1979 for a delimitation). We have not found or heard the species just south of the Amazon in the forests along the Manaus-Humaită road, but M MARTINS and M. GORDO (pers. comm.) Found specimens at the Rio Urucu about 100 km SSE of Coari. It has also been recorded from the Parque Nacional do Tapajós near Itaituba, southwest of Santarém, Pará (ZIMMERMAN & BOGART, 1984).

Discussion. — The new species possesses a number of characters new or unusual for the genus. In their definition, TRUEB & DUELLMAN (1971) (slightly altered by DUELLMAN & HOGOMOED, 1992) found Osteocephalus to have "vocal sacs paired, posterior, and when inflated protruding posteroventral or posterolateral to angles of jaws". Osteocephalus ophagus has a single median subgular vocal sac, a character shared with some frogs of the O. rodriguer! group (sensu AYARZAGUENA et al., 1992a) (AYARZAGUENA et al., 1992b). Also, it is similar to some frogs of that group in that the dorsal skin structure is not a reliable sexually dimorphic character (AYARZAGUENA et al., 1992a). Although males tend to be slightly more granular, there are smooth skinned males both alive and especially in preservative.

Tadpoles have two upper and three lower tooth rows like most lentic hylid larvae. In other species of Ostcocephadus, however, there are 2-3 upper and 5-8 lower tooth rows (HENLE, 1981; HERO, 1990). The reduction of denticle rows in O. oophagus may be an adaptation for oophagy as a special case of macrophagy. The tadpoles do not need to rasp their food as grazers. This might also be the reason for the absence of beak serrations. The reduction of denticle rows is known for many other arboreal tadpoles (LANNOO et al., 1987) and does not oppose its inclusion in the genus Ostcocephalus.

Despite the differences mentioned above, *O. oophagus* shares important characters with *O. taurinus*: the juvenile colorations and the colour of the iris in adults are identical. The frontoparietal flanges are present, though less conspicuous in *O. oophagus*. For these reasons the new species may be most closely related to *O. taurinus*.

Derivatio nominis. - The specific name oophagus is a compound of the Greek oon (egg) and phagein (to eat) and refers to the larval habit of eating conspecific eggs.

RESUMEN

Osteocephalus oophagus sp. nov. de Reserva Ducke, Manaus, Amazonas, Brasil, sepecie mediogrande del género, es semejante a O. taurinus pero más pequeña y con las piernas posteriores más cortas. En contraste a otros Osteocephalus, la piel dorsal no muestra dimorfismo escual y el saco vocal del macho es impar, mediano y subgular. El renacuajo es atipico para este gênero, porque vive en fitotélmata, alimentándose de huevos coespecíficos y teniendo una fórmula de dentículos labiales de 2(2)/3. La especie es conocida de las bajas Guianas y de la baja Amazonía.

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Reassessment of central Peruvian Telmatobiinae (genera Batrachophrynus and Telmatobius). I. Morphometry and classification

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The taxonomic status of the central Peruvian Telmatobilina is reassessed by analyzing the intraspecific variation of 18 morphometric measures among the currently recognized taxa (3 genera, 6 species, 9 subspecies). Cluster analysis, principal component analysis and discriminant analysis lead to the recognition of two genera (Batrachophynus and Telmatobius) including six species (Batrachophynus brack) and thout segregation in subspecies. Two diagnostic external features distinguish Batrachophynus probus species, another two characters are convergent adaptations distinguishing the stema-inhability acoupter from the lake-inhabiting one.

INTRODUCTION

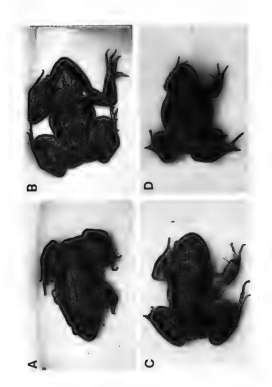
Leptodactylid frogs of the genus Telmatobius Wiegmann, 1835 inhabit the whole range of the Andes from Ecuador in the north to Chile/Argentina in the south (FROST, 1985; CEI, 1986). However, in the high-Andean regions of central Perú around Lake Junin (FJELDSA, 1983), two endemic Telmatohius-like species have been described as members of the genus Batrachophrynus Peters, 1873: the large lake-inhabiting Junin frog B. macrostomus and the stout stream-inhabiting B. brachydactylus. PETERS's (1873) distinction was based solely on the presence (Telmatobus) or absence (Batrachophrynus) of maxillary and prevomerine teeth. Yet, T. brevipalmatus, T. edentatus and T. intermedius lack maxillary teeth (LYNCH, 1971). Differences in several osteological characters other than teeth between Batrachophrynus and Telmatobus confirmed the generic distinction (LYNCH, 1978) and indicated an early separation of Batrachophrynus from the Telmatobius stock (CEI, 1986), LAURENT (1983) assigned B, brachydactylus to a monotypic third genus Lynchophrys based on morphometric differences from B. macrostomus. At present, this view on taxonomy is maintained, though independent support is absent (DUELLMAN, 1993; FROST, 1985; LAVILLA, 1988a). According to LAURENT'S opinion, Lynchophrys is more recently derived from the Telmatobius stock than is Batrachophrynus. Thus, the

phylogenetic relationship among the two species B. brachydactylus and B. macrostomus and of both with the genus Telmatobius are controversial.

The central Peruvian Tehmatobius presently include four stream-inhabiting species, T. brevirostris Vellard, 1955, T. carrillae Morales, 1988, T. Jelskii (Peters, 1873) and T. rumac Schmidt, 1954, whereas T. juminensus (Shreve, 1938) was an incorrectly classified Phrynopus (DTELLMAN, 1993). VELLARD (1955) proposed three subspecies for T. brevirostris, four for T. jelskii and two for T. rumac, although the significance of the morphometric differences between the proposed subspecies was not tested for, and the existence of intermediate individuals was stated. Field studies in the Mantaro Valley (Department of Junin, Perú) on T. jelskii raised doubts on the subspecific classification because morphs pertaming to different "subspecies" were found within the same population (Shixett, 1985, 1986, 1990).

Part of the taxonomic confusion in this group is probably due to morphological convergence among the species of each ecotype: the stream-inhabiting, semiaquatic frogs are stout and usually moderate-sized or small, whereas the lake-inhabiting, fully aquatic frogs are large (SINSCH, 1986, 1990). Therefore, phenotypic similarity between different species may reflect convergent adaptation to the same environmental factor rather than phylogenetic relationship. On the other hand, subtle differences among populations of the same species could result from direct responses to such factors as the amount of water flow in different streams. To test for environmental influences on morphological traits, we complemented our analysis of the two *Batrachophrymus* species and four stream-inhabiting *Telmatobius* species from central Perú with the lake-inhabiting Titicaca frog *Telmatobius* culeus (Garman, 1875) from southern Perú.

It is obvious that the validity of the central Peruvian taxa of Telmatobiinae needs to be confirmed by a thorough investigation. The definition of most taxa is exclusively based on morphological and morphometric characters (e.g. VELLARD, 1951, 1953, 1955) without an appropriate assessment of the intraspecific variation (TRUEB, 1979). Currently, the identification of most species is only possible by comparison with type specimens Consequently, our investigation evaluates VELLARD's type material in MHNSM (Museum of Natural History "Javier Prado", Lima, Perú) in comparison with the preserved specimens in URP (Museum of Natural History of the Ricardo Palma University, Lima, Perú) and with specimens collected in the field in 1992. We use cluster analyses to detect intraspecific morphological traits within the data sets of phenotypes described by morphometric characters. At the species level we widely follow the procedures used by WIENS (1993) in his recent taxonomic revision of the Telmatobiinae from northern Perú. Principal component and discriminant analyses are applied to quantify the differentiation among the seven nominal species. Finally, the differences among the genera and the ecotypes (stream- versus lake-inhabitants) are surveyed in discriminant analyses. The aims of our study are to provide an objective basis for the identification of central Peruvian Telmatobunae and to distinguish convergent morphological traits from morphometric features which characterize species and/or genera.



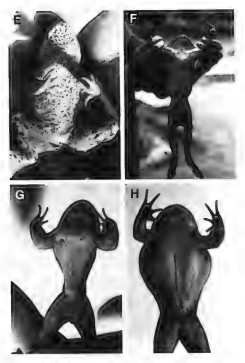


Fig. 1 Batrachophrynus brachydaetylus (A, dorsal view; E, ventral view); B. macrostomus (B, F); Telmatohus jelsku (C, G); T. rumac (D, H)

MATERIAL AND METHODS

The material examined included a total of 280 adult frogs pertaining to the seven nominal species Batrachophrymus brachydactylus (figs. 1A, 1E), B. macrostomus (figs. 1B, 1F), Telmatobius brevirostris, T. carrilae, T. culeus, T jelskii (figs. 1C, 1G) and T. rimac (figs. 1D, 1H). The frogs had been collected at 32 localities (fig. 2). The assignment of specimens to subspecies and to localities is listed in Appendix 1, the number of individuals studied and the sex-specific size are given in Table 1.

Standard morphometric measurements (nearest 0.1 mm) include: (1) snout-vent length (SVL); (2) height of body at the pectoral girdle (BH); (3) maximum width of head (HWID); (4) eye diameter (EYE); (5) interorbital distance (IOD); (6) eye-nostril distance, from anterior margin of eye to posterior edge of naris (ENOSE); (7) distance between the eye and the tip of the snout (ESNOUT); (8) humerus length (HUML); (9) radioultar length (RADL); (10) hand length (HNDL); (11) length of the third finger (FG3L); (12) femur length (FEML); (13) tibia length (TIBL); (14) foot length, from union with tibia to the tip of fourth toe (FOOTL); (15) length of first toe (TOEL); (16) length of fourth toe (TOE4L); (17) length of callus internus (CLL); (18) maximum length of toe web (WEBL).

Multivariate analyses were performed on log₁₀-transformed data (BooksTEIN et al., 1985) and morphometric ratios The empiric measurements were transformed to ratios (range: 0-1) by calculating measures relative to SVL (SCHNEIDER et al., 1992, 1993). Moreover, two indices were used for further analysis: CIL/TOE1L and FEML/TIBL. Table II gives the means (and corresponding standard deviation) of these 19 relative measures for each species.

Due to the low number of individuals assigned to type material, we analysed the morphometric similarity between individuals in a cluster analysis using hierarchical grouping of the SVL-standardized phenotypes (WARD, 1963). This procedure subsequently reduces the number of groups by joining that specimen to another one or to a cluster which organates the lowest error sum of square. The result is a dendrogram based on pheneuc similarity. As the proposals of subspecies by VELLARD were exclusively based on external morphology, valid subspecific taxa are expected to form homogeneous clusters.

At the species level, sets of the log_{in}-transformed data were subjected to principal component analysis to explore the morphometric variability independent of taxonomic assignment. Principal components (PC) are linear combinations of the measured variables, uncorrelated with each other and explaining the maximum amount of variation. The first principal component (PC) of morphometric data generally describes differences in size, but size effects may be present in subsequent principal components (HUMPHIRES et al., 1981). Techniques such as shearing have been developed to correct PC2 and PC3 for possible size effects (BOOKSTEIN 1987). Moreover, we applied canonical discriminant still persist (ROHLF & BOOKSTEIN 1987). Moreover, we applied canonical discriminant analysis to distinguish between the taxonomic groups delimited a priori. The resulting discriminant functions (CAN = canonical variables) are linear combinations of the measured variables that maximize the differences between the groups. Discriminant

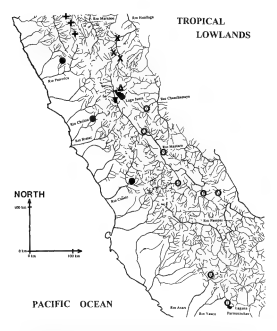


Fig. 2. – Distribution of the central Peruvan Telmatohunae. Butrachophryms brachydacrybac (inverted triangle); B macrostomus (open triangle), Telmatohun brewrostra (×), T carrillae (+), T jelskil (icircles), T rimac (dots) Localities are approximated from distances by roads, multiple localities in close proximity are represented by a single symbol. The main Andean river systems are indicated.

	B brach	edactylus	B macr	ostonus	T brev	nrostris	T cu	miliae	To	uleus	T J	elska	r,	ттас
Character	Males	Females	Males	Pernales	Males	Females	Males	Females	Males	Females	Malos	Permales	Males	Females
	N = 29	$N_1 = 24$	N = 6	N = 7	N = 2	$N \approx 3$	N = 32	N = 21	N = 19	N = 23	N = 37	N = 35	N = 26	N = 16
S۱L	56.5 ± 6.3	573 ± 56 449 694		133.0 ± 17.2 117.7 170.3	577 570 583	59 4 50 4 76 3	43 1 + 4 7 35 3 - 52 4	479±37 412-559		90 5 ± 23 5 50 7 ± 19 0		54 8 ± 5 4	49.8 ± 4.3	58 7 ± .0 9
BH	11.1 ± 7.2	106+29	230±46	234 2 4 8	16.4	17.1	10.2 ± 1.5	10.9 ± 1.2	570-1164 262±47	279 ± 92	176±14	43 9 71 2 13 9 ± 1 6	42 0 57 1 11 8 ± 1 6	47 0 86.9 14 3 ± 3 4
HWID	71-150	74-179 174±16	18 2 - 29 9 50 8 ± 4 7	183-330 554 ± 93	152-175 187	13 0 - 23 2 21 8	60-140 133±15	83·127 141+12	16.3 350 369 + 77	130 412 36.6 + 114	102-159	11 2 - 16 4 20 2 + 2 3	86-147 166+14	108-235 200+50
EYE	150-243 49+07	15 2 20 5 4 7 ± 0 6	461-574 82±14	464-753 76+07	17 4 · 20 0 5 8	17 0 30.1	104-162 38±07	115-170 39+05	213-519 70±10	168-517 68±12	157-233 53±08	151-257 55±08	138-183 54+0.6	144-316
IOD	32-68 137 ± 10	37 62	60 100 308 + 24	66 88	58-58	51-83	27-61	30-56	43-89	45-83	42-70	35-70	4.6 69	48 8.1
	117-156	117-152	277-339	32 0 ± 4 8 28 4 - 42 5	159-169	16 4 14 4 - 20 4	108±11 74-134	110±08 91-126	22 4 ± 3 4 14 2 - 27 1	216±54 120 287	11 4 20 2	148±14 110 L77	142±1, 126 161	159±28 131-238
ENOSE	87±13 66-139	83±06 67-94	184±14 16.2-198	19 2 ± 2 6 16 8 - 24 8	95 90-100	100 83-133	6.9 ± 1 3 5 5 - 12 9	68±05 56.77	124±19 86 156	127±28 80 167	90±07 80-107	91±08 75-110	86±06 76 98	96±19 79-153
ESNOUT	117±10 94-138	115 ± 0.9 96-134	274±19 249-297	28 3 ± 3 7 25 0 - 36 2	13.7 12.5 - 14.8	14 1 11 7 - 18 6	94 ± 08 73 113	96±05 89 L08	194 ± 28 127-250	189±43 117-239	129±09 114-153	130±10 108-146	123±08 106-137	136±25 115-209
HUML	154 ± 23 110 - 195	142 ± 17 105 167	49 3 ± 2 5 44 8 - 51 7	48 8 ± 7 0 42 2 - 62 8	17.4	173	10 8 ± 1 6	99±16 77-144	294±58 177-377	272±76	15.6 ± 2.1	154±24	144 ± 10	152 ± 33
RADL	115 ± 12	104 ± 12	342 ± 31	346±43	130	13.3	82-141 94±12	10 L ± L 2	216 ± 40	137-385 203±58	113-198 124±14	10 0 20 2 12 2 + 1 4	120-168 109±10	10 5 - 23 8 12 4 ± 2 1
HNDL.	97-144 113±11	82 127 111 ± 10	29 7 - 37 3 38 5 ± 2 3	30 8 - 42 0 39 9 ± 5 1	12 8 13 L 15 0	10 5 - 17 7 16 2	74-135 97+15	84-127 98±12	14 0 - 28 7 24 0 ± 4 3	10 1 - 27 3 23.4 ± 6 1	97-155 136±10	87-149 141±10	88 129 130±11	96 176 148±30
FG3L	91-139 63+07	94-131 64+09	361-425 251±16	36.1 - 51.0 24.5 + 3.1	148-152 94	14 6 - 19 0 10 9	63-129 57±09	78-122 59±10	15 2 - 31 0 14 6 ± 3 0	129 337 146±43	116-163 88±13	123-173 90±09	10.6 49 83±11	12.0 24.0 96 ± 2.6
FEML	50-80 255+34	50-83 243+32	232-276 623+17	21.4 - 30.4 63.5 + 4.0	94-94	94-139 284	42-76	38-81 206±14	77-189	6.0 21 J 39.8 + 10.5	6.9 113	73-106	6.0 10.6 23.8 ± 1.5	70-180 270+47
	20 4 - 34 8	19 5 - 32 7	59 5 - 66 3	578-689	24 4 - 25 9	250-315	14 4 24 Z	167-233	26.8 - 53.6	183-524	187-290	180 313	201-280	210-374
TIBL	$\begin{array}{r} 23 \ 6 \ \pm \ 2 \ 8 \\ 18 \ 6 \ \ 30 \ 3 \end{array}$	$\begin{array}{c} 228 \pm 26\\ 189 & 294 \end{array}$	56 4 ± 4 5 50 9 62 5	59 2 ± 6 5 53 4 - 73 0	273 258288	28 1 24.1 - 35 0	20 3 ± 2 2 16 4 - 24 5	209±18 180 256	27 1 - 56 8	41.4 + 10.9 20.4 - 57.5	21 4 - 26 8	24 2 + 1.6 20 5 - 27 0	25 1 ± 1 5 21 0 - 28 3	27 6 ± 4 3 21 4 - 36 8
FOOTL	$\frac{361 \pm 44}{310 \cdot 494}$	365+39 302-451	925±46 861-983	981±67 910-1123	41 3 40 3 - 42 2	44 6 38 3 - 53 7	316+38 255-405	32 9 ± 3 0 29 3 · 41 0	71 0 ± 12 9 46.2 - 86 4	67 2 ± 17 1 35 3 90 0	379 ± 30 290 - 423	379±35 250×422	381±28 328-427	438±70 372-624
TOEIL	49±06	\$1±06 37.65	159±1.6 136 183	175 ± 24 152-326	64 54 73	66 55-86	44 ± 05 36 55	50 ± 07 37 60	10 2 ± 2 2 5 8 - 13.4	98±30 50-144	49±05 39-58	48±07 37-59	58±06 43-67	68±12 54-103
TOE41	34 0 ± 2 7	23.4 ± 2.3	588±29	60 8 ± 3 2	27 3	29.2	206+23	22 0 ± 1 9	477 + 88	456 + 121	24.4 ± 2.2	251 ± 28	258±22	299 ± 49
CiL	206 295 23±05	23±05	55 6 62 9 4 6 ± 0 7	56 0 65 2 4 5 ± 0 9	36	265-336 34	176 - 265 22 ± 04	194 - 267 22±03	29 2 - 57 7 4 5 ± 1 0	251-677 43±12	167-280 24±06	136-290 27±06	21 1 · 28 6 2 6 ± 0 3	24 4 41 7 29 ± 06
WEBL	.635 74±15	16-38 69±14	37.52 292±37	34-55 271±39	33-38 76	32-36	14·28 68±17	17-31 78±10	27 65	26 6.5 181±65	11 32 98 ± 38	1543 104±38	20 31 97±45	21 4.4
	4 2 - 11 3	43 96	24.5 - 34.6	22 2 - 34 5	4 2 10 9	32.83	39 1.0	62 101	95-252	73-274	44 .60	40 175	38-178	54-233

Table 1 Morphometric data for Batrachophrinus brachidactilus, B macrostomus Telmatobius brevirostris T carrilloe T culeus, T jelsku and T rimac. The first line is mean ± 1 SD; second line is range. All values are in millimeters, see text for abbreviations of variables.

Ratio	B. brachydactylus	B. macrostomus	T brevirostris	T. carrillae	T. culeus	T. jelsku	T. rimac
Hano	N = 53	N = 13	N = 5	N = 53	N = 42	N = 72	N = 42
BH/SVL	0.194 ± 0.047	0.176 ± 0.025	0 285 ± 0 023	0.231 ± 0 020	0 292 ± 0 032	0.252 ± 0 028	0.240 ± 0.029
HWID/SVL	0 314 ± 0 026	0.404 ± 0.024	0.347 ± 0.034	0.302 ± 0.021	0.397 ± 0.037	0.364 ± 0.026	0.337 ± 0.026
EYE/SVL	0.085 ± 0.014	0 060 ± 0 007	0 103 ± 0.004	0.085 ± 0.012	0 076 ± 0.009	0.100 ± 0.015	0 106 ± 0 012
IOD/SVL	0.240 ± 0.021	0.238 ± 0.010	0 281 ± 0 011	0.242 ± 0.019	0.240 ± 0.014	0.270 ± 0.026	0.282 ± 0.022
ENOSE/SVL	0.151 ± 0.015	0 143 ± 0.005	0.167 ± 0.006	0.152 ± 0.022	0.138 ± 0.012	0.167 ± 0.014	0 170 ± 0 013
ESNOUT/SVL	0.206 ± 0.016	0.212 ± 0.007	0 236 ± 0 013	0.210 ± 0.014	0 211 ± 0 018	0.238 ± 0 015	0 243 ± 0.016
HUML/SVL	0.262 ± 0.034	0 373 ± 0.020	0.299 ± 0.027	0.233 ± 0.037	0.306 ± 0.025	0.285 ± 0.043	0.280 ± 0.028
RADL/SVL	0.195 ± 0.022	0.262 ± 0.017	0.227 ± 0.014	0.215 ± 0.017	0 227 ± 0 015	0.226 ± 0 023	0 218 ± 0.021
HNDL/SVL	0.199 ± 0.021	0.299 ± 0.017	0.270 ± 0.020	0.217 ± 0.027	0.257 ± 0.017	0.255 ± 0.023	0.260 ± 0.022
FG3L/SVL	0.113 ± 0 015	0.189 ± 0 017	0.176 ± 0.012	0.128 ± 0.017	0 158 ± 0 019	0.163 ± 0.025	'0.166 ± 0.024
FEML/SVL	0 440 ± 0 038	0.481 ± 0.039	0.468 ± 0.062	0 445 ± 0.037	0.452 ± 0.032	0.453 ± 0.043	0.475 ± 0.039
TIBL/SVL	0 410 ± 0 035	0.441 ± 0 019	0.475 ± 0.021	0.457 ± 0.027	0 467 ± 0 034	0.448 ± 0.033	0.495 ± 0.037
FOOTL/SVL	0 640 ± 0 047	0.729 ± 0.039	0.741 ± 0.051	0 713 ± 0.042	0.751 ± 0.062	0.695 ± 0.070	0.763 ± 0.044
TOE1L/SVL	0 088 ± 0 010	0.127 ± 0.010	0.110 ± 0.011	0.104 ± 0.013	0 108 ± 0.013	0.089 ± 0.011	0.117 ± 0.012
TOE4L/SVL	0.418 ± 0.030	0.458 ± 0.033	0.489 ± 0.047	0 469 ± 0.029	0.506 ± 0.042	0.454 ± 0.041	0.519 ± 0.047
CIL/SVL	0 041 ± 0 009	0.035 ± 0 006	0 060 ± 0.009	0.049 ± 0.006	0 047 ± 0.007	0.047 ± 0.011	0.051 ± 0 008
WEBL/SVL	0.126 ± 0.026	0.214 ± 0 027	0 101 ± 0.053	0 159 ± 0.027	0.198 ± 0 036	0.187 ± 0.072	0.191 ± 0.087
CIL/TOE1L	0.472 ± 0.107	0.275 ± 0.050	0.555 ± 0.123	0 484 ± 0.092	0.440 ± 0.066	0.536 ± 0.144	0.442 ± 0 056
FEML/TIBL	1.072 ± 0.074	1 092 ± 0 084	1 002 ± 0.090	0.975 ± 0.075	0.972 ± 0.074	1.013 ± 0.080	0.960 ± 0 058

Table II. - Ratios of morphometric data for Batrachophronus brachodactilus, B. macrostomus, Teimatobius brevirostris, T. carrillae, T. culeus, T. jelskii and T. rimac. Data are given as mean ± 1 SD. See text for abbreviations of variables.

functions were derived from the log₁₀-transformed data and from the ratios. The degree of separation of taxa was almost identical in both analyses and therefore we present the results using the log₁₀-transformed data only (analogous to Wiens, 1993).

All calculations were performed on a PC using the FORTRAN77 program CLUSTER and the program package STATGRAPHICS, version 5.5.

RESULTS

INTRASPECIFIC VARIATION

Among the seven nominal species studied in this paper, four are thought to segregate in subspecies: Tehnatolius hervirostrist, T. culeus, T. jetkii and T. rimae (VELLARD, 1951, 1953, 1955). We approached the problem of morphometric variation within a nominal species by applying cluster analysis on the phenotypes of all conspecific individuals available, including those which VELLARD assigned as type material for the proposed subspecies. Phenotypes are described by 19 SVL-standardized morphometric ratios to minimize size effects.

The phenograms of all species (including those without subspecific segregation) revealed existence of different intraspecific morphological traits documented by groups of specimens which joined to the same cluster with an error sum of squares of less than 0.1 (figs. 3-6, data on *Batrachophrynus* and *T. carrillae* not shown). We treat these groups of remarkably similar specimens as "morphs", without intending a taxonomic implication. Generally, intraspecific morphs did not reflect polymorphism related to sex or size.

Telmatobius brevirostris Vellard, 1955

VELLARD (1955) recognized three subspecies: Telmatobus b. brevitostris, T. b. parvalus and T. b. punctatus. Unfortunately, we only found in the MHNSM collection two specimens of the first two subspecies, and one of the third. The original descriptions were based on only three adults of T. b. brevitostris, sux of T b parvalus and two of T. b. punctatus – a prohibitively small sample size to define reliable subspecific taxa. The phenogram (fig. 3) of the five specimens available for analysis shows that: (1) the two type specimens of T. b. brevitosits from Chasqui join different clusters; (2) the morphometrically most similar specimens are one T. b. brevitostris individual and the T. b. punctatus male from Stant Maria del Valle; (3) the two individuals of T. b. parvalus from Caina are very alike, but form together with one T. b. brevitostris a main cluster with a error sum of squares far below the level of morph distinction. All specimens join to one group at an error sum of squares of ontpoly 0,146. Nether the association of the five phenotypes to groups nor the degree of morphological differentiation between them support a subspecific segregation within the nominal species T. brevitostris.

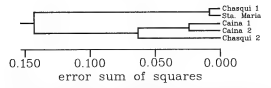


Fig 3 — Phenogram of morphometric similarity between adult Tehnatohus brevitaritis which VELLARD (1955) assigned to different subspecies Each specimen is identified by the locality of collection and an individual number Similarity is based on 19 ratios and computed by hierarchical grouping in a cluster analys. T. b. brevitostris: Chasqui, T b. parvulus: Cana, T. b. puertains. Santa Mara del Valie.

Telmatobius culeus (Garman, 1875)

VELLARD (1953) recognized six subspecies: Telmatobius c. culeus, T. c dispar (redescribed by LAVILLA, 1988b), T. c. escomeli, T. c exsul, T. c. fluviailis and T. c lacustris. The phenogram (lig. 4) calculated for 42 adults assigned as type material (MHNSM) reveals that: (1) there exist five morphs within the nominal species; (2) these morphs do not coincide with any of the subspecies proposed by VELLARD; (3) different morphs occur at the same locality. Joining all specimens to one group causes an error sum of square of 0.453, that is three times greater than in T brevirostris. Again, from the morphometric point of view there is no evidence that the morphological traits within T. culeus agree with the proposed subspecific differentiation.

Telmatobius jelskii (Peters, 1873)

VELLARD (1955) recognized four subspecies. Telmatobius j jelskii, T. j. bufo(redescribed by LAVILA, 1988b), T. j. longitarsis and <math>T. j walker! The phenogram (fig. 5) of 72 adults including VELLARD's type specimes (MHNSM) shows that (1) there exist seven morphs within the nominal species; (2) these morphs do not coincide with any of the subspecies proposed by VELLARD; (3) different morphs inhabit the same locality. Joining all specimes to one group causes an error sum of squares of 1.44M, that is three times greater than in T. culcus or in T. rimac and ten times greater than in T brevirositis. Nevertheless, the occurrence of different morphs at all sites – though in different frequencies – does not support the validity of the four subspeces proposed by VELLARD.

Telmatobius rimac Schmidt, 1954

VELLARD (1955) recognized two subspectes: *Telmatobius r. rimac* and *T. r. merahonalis*. The phenogram (fig. 6) of 42 adults including VELLARD's type spectmens shows that: (1) there are five morphs; (2) the type material from Ocros and Tupe forms one of two

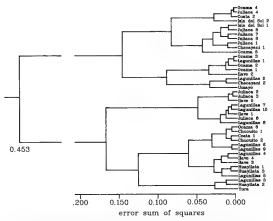


Fig 4 – Phenogram of morphometric similarity between adult *Telmutobius culeus* which VELLARD (1953) assigned to different subspecies. Presentation of data analogous to that in fig. 3. *T. c. culeus* Ocama, liave, Isla del Sol, *T. c. dispar*: Coata, Juliaca; *T. c. escometr.* Lagunillas; *T. c. exuit.* Yura; *T. c. fluvantis.* Chucuito, Huxgilata, Ilave; *T. c. lacustris.* Checayani, Umayo

main clusters including two mixed morphs; (3) the more recently collected material from Obrojillo and Huaytara (situated at the same distance to Ocros in the north and to Tupe in the south) forms the second main cluster and segregates into three more morphs; (4) only one individual (Obrojillo 3) directly joins the cluster of the type specimens. This invexpected structure of the phenogram does not support the validity of the proposed subspecies, but demonstrates suspicious differences between the external morphology of T. *rinuc* which were collected forty years ago and those presently inhabiting the streams of the Paofic Slope of the Andes.

INTERSPECIFIC VARIATION

In order to compare the intraspecific variation with the morphometric differences among the nominal species, we performed principal component analysis and discriminant analysis on two groups of geographically neighbouring species: (1) the northern Tel-

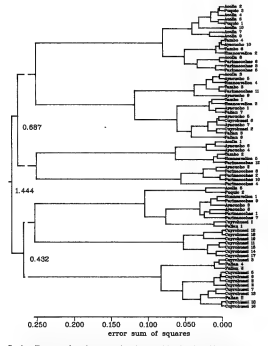


Fig. 5 Phenogram of morphometric similarity between adult Telmatohus jelskii which VELLARD (1955) assigned to different subspecies. Presentation of data analogous to that in fig. 3 T. j. jelskii Acolla; T j bufo Tambo; T. j. longitarist: Huancavelica, Puquio; T. j. walkeri Ayacucho. Unclassified T. jelskii were collected in Palian, Cuyrohuasi and Parinacochas

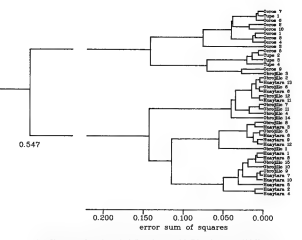


Fig. 6. – Phenogram of morphometric similarly between adult *Telmatobus rumae* which VELLABD (1955) assigned to different subspecies. Presentation of data analogous to that in fig. 3 *T. r. rumae*: Ocros, *T. r. meridionalis* Tupe. Unclassified *T rimae* were collected in Obropillo and Huaytare.

matobinae including Batrachophrynus brachydactylus, B. macrostomus, Telmatobius brevirostris and T. carrillae; (2) the southern Telmatobius species T. culeus, T. jelski and T. rumac. The main reason for the subdivision of the complete data set on phenotypes into groups of three and four species, respectively, was to reduce the number of significant canonical variables to two, which permits the distinction of these taxa in two-dimensional scatter plots (figs. 7-8). Here, we present only the results based on \log_{10} -transformed data because the separation of taxa by discriminant analysis was almost identical in data sets of 18 log₁₀-transformed distances and in those consisting of 19 morphometric ratios.

Generally, the interspecific differences in size (PC1) by far exceeded those in shape (PC2, PC3). The size effects on PC2 and PC3 appeared to be small, because shearing did not notably improve the separation of taxa. Discriminant analysis led to an almost optimal separation of species by combining differences in size and shape.

In the northern group of central Peruvian Telmatobinae, the first three principal components explained 95.0 % of the total variance. PCI distinguishes the large B. macrostomus from the smaller three species. The plot of PC2 and PC3 scores (fig. 7A) shows a wide overlap between B. macrostomus, B. brachydactylus and T. carrillae, whereas PC3 distinguishes four of the five T. brevnostrus from the other species An almost complete separation of the four taxa was obtained by discriminant analysis, only 2 out of 53 B. brachydactylus were confounded with T. carrillae (fig. 7B, Table III). The separation of taxa is mainly based on size (SVL) and interorbital distance (IOD) in CAN1, and on size (SVL), head shape (HWID, EYE, IOD) and humerus length (HUML) in CAN2.

In the southern group of Telmatobus species the first three principal components accounted for 92.9 % of the total variance. PCI distinguishes T. culeus from the other two species. The plot of principal component scores (fig. 8A) shows a considerable separation of T. yelskii from T. rimac based on PC3, but a complete overlap of T. culeus with both other species. An almost complete separation of the three taxa was obtained by canonical discriminant analysis, only 1 out of 42 T. culeus was confounded with T. yelskii, and 1 out of 72 T. jelskii with T. rimac (fig. 8B, Table IV). The separation of species is mainly based on size (SVL, BH) and head shape (ENOSE, ESNOUT) in CAN1, and on size (SVL), head shape (HWID, ESNOUT) and extremity length (HANDL, TIBL) in CAN2.

INTERGENERIC VARIATION

The next step of analysis concerns the morphometric features of each genus and the possibility to identify genus-specific morphometric characters or ratuos. The data of the two *Batrachophrymus* species form one group, those of the five *Telmatobius* taxa the other. Again, \log_{10} -tranformed data and ratios provided the same degree of group separation A highly significant discriminant function was obtained which correctly classifies 80 % of the *Batrachophrymus* and 96 % of the *Telmatobius* (\log_{10} -transformed data, Table V). Distinctive morphometric ratios (ANOVA, P < 0.01) were BH/SVL (fig. 9A) and FEML/TIBL (fig. 9B).

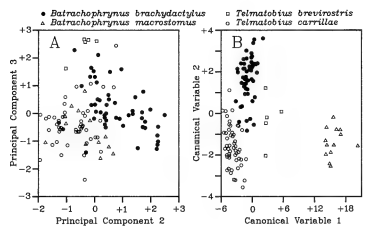


Fig 7. Plot of (A) principal component scores and (B) discriminant function scores of the four northern telinatobine species: Bartachophyrum brachopharylus, B. macrossonus, Telinaiobus brevirositris and T. carrillae. Discriminant functions and classification success are given in Table III.

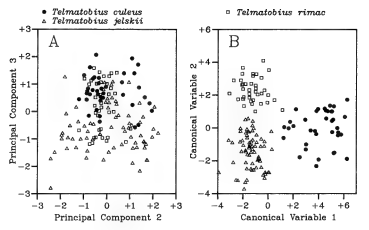


Fig. 8 — Plot of (A) principal component scores and (B) discriminant function scores of the three southern telmatobium species' *Telmatobius pelskin*, *T. rimac* and *T. culeus*. Discriminant functions and classification success are given in Table 1V

Table III - Discriminant functions to distinguish among Batrachophrynus macrosiomus, B. brachydactylus, Telmatobus brevrostrs and T. carrillae based on 18 logio transformed morphometric characters.

A Statistical significance

Eigenv	alue	Canonical correlation	Wilks Lambda	Chi-squared	Degrees of freedom	р
32 1	3	0.985	0.0042	613 5	54	< 0.00001
3 4		0 842	0.1383	221.5	34	< 0.00001
1.1		0.725	0.4741	583 6	16	< 0.00001

B. Unstandardized discriminant function coefficients

Character		Coefficients	
Character	CAN I	CAN 2	CAN 3
SVL	- 11 64	18 89	- 6.59
BH	- 3.60	- 0.10	4.81
HWID	2 91	17 56	- 9.19
EYE	- 6.78	10 25	2 81
IOD	25 39	- 12.21	14.51
ENOSE	- 4.42	- 4.49	- 10.26
ESNOUT	6.79	- 7 75	7 84
HUML	1 64	11.01	0.92
RADL	5 77	- 7 09	- 3.67
HNDL	6 99	- 5.13	- 1.41
FG3L	6 92	- 1 94	6.82
FEML	9 23	6 85	- 4.27
TIBL	- 5.42	- 8 20	6.23
FOOTL	- 8 38	1.21	- 4.85
TOE11	4.57	- 7,16	1.60
TOE4L	2 50	- 8 12	7.13
CIL	- 2.14	- 0.50	3.27
WEBL	- 1 62	- 3.84	- 9.81
Constant	- 25.08	- 13.17	0.75

C. Classification success

		Predicted group					
Actual group	Batrachophrynus	Batrachophrynus	Telmatobuo	Telmatobuci			
	brachydactylus	macrostomus	brevirostris	carrillae			
B. brachydactylus	51 (96%)	0	0	2 (4%)			
B. macrostomus	0	13 (100%)	0	0			
T. brevirostris	0	0	5 (100%)	0			
T. carrillae	0	0	0	53 (100%)			

D Group centroids

Species	CAN 1	CAN 2	CAN 3
B. brachvdactvlus	- 0.71	1.75	- 0.16
B macrostomus	15.59	- 0 98	- 0.58
T. brevirostris	- 3 41	- 1 49	- 0.17
T. carrillae	3.17	- 0.24	5.02

Table IV. - Discriminant functions to distinguish among *Telmatobius culeus*, *T. jelskii* and *T. rimac* based on 18 log₁₀ transformed morphometric characters.

	Α.	Statistical	significance
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Eigenvalue	Canonical correlation	Wilks Lambda	Chi-squared	Degrees of freedom	Р
5 47	0.919	0.0416	440.3	36	< 0.00001
2 71	0.855	0.2694	181.7	17	< 0.00001

B Unstandardized discriminant function coefficients

Chanadar	Coeff	icients
Character	CAN 1	CAN 2
SVL BH HWID EYE IOD ENOSE ESNOUT HUML RADL HADL FG3L FG3L FG3L FOOTL TOBL TOBL TOBL CIL	- 0.53 11,13 0,72 -5,65 -2,364 -11,85 7,22 -3,76 3,76 -3,76 -5,56 -6,56 -6,56 -5,17 -2,28 -3,76 -3,77 -6,56 -3,77 -6,56 -3,77 -6,56 -3,77 -2,28 -3,76 -3,76 -3,76 -3,76 -3,64 -3,76 -3,76 -3,64 -3,76 -3,76 -3,76 -3,64 -3,76 -3,75 -3,76 -3,77 -3,76 -3,77 -3,76 -3,77 -3,77 -3,76 -3,77 -3,77 -3,76 -3,77 -3	$\begin{array}{c} 10\ 88\\ -0.17\\ -8.52\\ 3.52\\ 9.22\\ -0.28\\ -0.28\\ -1.20\\ -8.32\\ -10.22\\ -4.33\\ -0.34\\ 15.38\\ -10.38\\ -10.38\\ -10.38\\ -10.38\\ -10.38\\ -10.38\\ -10.38\\ -10.38\\ -10.38\\ -10.38\\ -10.38\\ -10.38\\ -10.25\\ -10.$
WEBL Constant	0.55 - 10.60	- 0 48 1.19

C. Classification success

	Predicted group		
Actual group	Telmatobius culeus	Telmatobius jelskii	Telmatobius rimac
Telmatobius culeus Telmatobius jelskii Telmatobius rimac	41 (98%) 0 0	1 (2%) 71 (99%) 0	0 1 (1%) 42 (100%)

D. Group centroids

Species	CAN 1	CAN 2
Telmatobius culeus	4.05	- 0 06
Telmatobius jelsku	- 1.37	- 1.42
Telmatobius rimac	- 1.25	2 46

Table V. Discriminant functions to distinguish the genera Batrachophrynus and Telmatobius based on 18 log₁₀ transformed morphometric characters.

A. Statistical significance

	Eigenvalue	Canonical correlation	Wilks Lambda	Chi-squared	Degrees of freedom	Р
ł	1.63	0.788	0.3794	254.9	18	< 0.00001

B. Unstandardized discriminant function coefficients

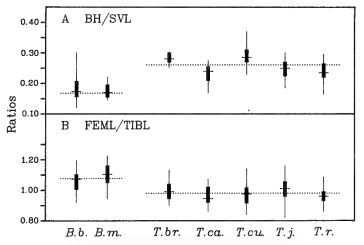
0	Coefficients	
Character	CAN 1	
SVL	- 19.89	
BH	8.83	
HWID	0.90	
EYE	- 0.77	
IOD	- 3.75	
ENOSE	- 10.09	
ESNOUT	11.96	
HUML	- 5.36	
RADL	- 0.61	
HNDL	6.05	
FG3L	4.03	
FEML	- 9.71	
TIBL	13.01	
FOOTL	- 3.93	
TOELL	- 2.39	
TOE4L	10.08	
CIL	- 0.23	
WEBL	0.56	
Constant	13.78	
Constant	13.78	

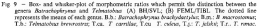
C. Classification success

	Predicted group	
Actual group	Batrachophrynus	Telmatobius
Batrachophrynus Telmatobius	53 (80%) 8 (4%)	13 (20%) 200 (96%)

D. Group centroids

Genus	CAN I
Batrachophrynus	- 2 26
Telmatobius	0.72





VARIATION BETWEEN ECOTYPES

Finally, we examined the relationship of morphological traits caused by the convergent adaptation to a specific type of habitat (stream/lake) with the phylogenetic relationships between the seven taxa. For this purpose, we pooled the data of *B.* macrostomus and *T. culeus* forming the lake group and compared them with the stream group formed by the remaining species. Again, there was no difference between the separatuon of groups based on log₄₀-transformed data and that based on ratios. The highly significant discriminant function correctly classifies 100 % of the stream-inhabitants and 2% so fith lake-inhabitants (log₄₀-transformed data, Table VI). Distinctive morphometric ratios (ANOVA, P < 0.01) between lake- and stream-inhabitants are HWID/SVL (fig 10A) and EYE/SVL (fig. 10B). Moreover, SVL of adult lake-inhabitants is considerably larger than that of stream-inhabitants (Table I).

DISCUSSION

The general similarity in size and shape of the riparian central Peruvian Tehnatobiinae does not facilitate a reliable and objective definition of taxa. TRUEB (1979) stated that most taxonomic descriptions of *Telmatobius* are inadequate because many proposed diagnostic characters are so subjective that an identification without comparison with type material is almost impossible. A recent study on the *Telmatobius* species of northern Perù used for the first time multivariate statistics on morphometric data to obtain objective criteria for the classification of telmatobiine frogs (Wiens, 1993). We followed this approach and successfully apphed cluster, principal component and discriminant analyses to assess the morphometric variation within and among the central Peruvian taxa of Telmatobiinae. Consequently, we offer discriminant functions based on external characters which permit the identification of adults with a very low rate of erroneus classifications. Moreover, we identified convergent morphological traits distinguishing riparian and lake-inhabiting telmatobiine frogs which in the future should be avoided for taxonomic conclusions.

REASSESSMENT OF THE SUBSPECIFIC TAXA

Within wide ranging species such as *T. jelskii* and *T. rimac* we are confronted with the problem of interpopulational morphological differentiation, due to the partial geographical isolation between different hydrographic systems and valleys. VELLARO (1951, 1953, 1955) attempted to solve this problem by naming more than half of the populations studied up to 1955 at the subspecific level. However, the existence of a complete series of intermediate specimes (VELLARD, 1955) between all subspecies emphasizes the more or less arbitrary nature of their definition. Nevertheless, due to VELLARD's proposal, the segregation of *T. brevirostris*, *T. culeus*, *T. jelskii* and *T. rimac* into a total of fifteen (!) subspecies is still recognized (FROST, 1985).

Our attempt to deal with interpopulational variability in taxonomic terms is based on the hierarchical grouping of individual phenotypes according to their similarity in external Table VI. - Discriminant function to distinguish between the stream- and lakeinhabiling species of the genera *Batrachophrynus* and *Telmatobius* based on 18 log₁₀ transformed morphometric characters.

A. Statistical significance

Eigenvalue	Canonical correlation	Wilks Lambda	Chi-squared	Degrees of freedom	Р
4.58	0.906	0.1792	452.1	18	< 0 00001

B. Unstandardized discriminant function coefficients

Character	Coefficients		
Character	CAN I		
SVL	3.21		
BH	4.72		
HWID	8.12		
EYE	- 4.01		
IOD	- 5.74		
ENOSE	- 8.33		
ESNOUT	0.77		
HUML	6.98		
RADL	~ 1.30		
HNDL	4.85		
FG3L	- 6.24		
FEML	- 3.29		
TIBL	- 3.84		
FOOTL	7.42		
TOEIL	3.14		
TOE4L	7.58		
CIL	0 24		
WEBL	0.87		
Constant	16.34		

C. Classification success

A studies out	Predicted group		
Actual group	Stream-inhabitant	Lake-inhabitant	
Stream-inhabitant Lake-inhabitant	224 (100%) 4 (8%)	0 46 (92%)	

D. Group centroids

Genus	CAN 1
Stream-inhabitant	- 1.01
Lake-inhabitant	4.51

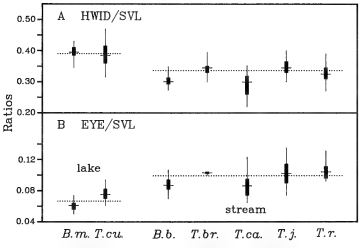


Fig 10 – Box- and whisker-plot of morphometric ratios which permit the distinction between inhabitants of lakes and of streams (A) HWID/SVL; (B) EYE/SVL. Presentation of data analogous to fig. 9.

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morphology To avoid major biasses due to size, the phenotypes are described by 19 unweighted morphometric ratios. The phenotypic grouping in a cluster analysis permits an objective assessment of similarity between conspecific individuals, even if the sample size is small as usually for type material. Principal component and discriminant analyses, in contrast, require about 20 cases per predictor which imply sample sizes exceeding by far the numbers of type specimens.

VELLARD's proposal of subspecies is exclusively based on differences in the external morphology of specimens from different localitus. Therefore, if the proposed subspeces were valid, the phenograms of conspecific individuals collected from different localities (= reproductively isolated populations) should show the following structure: (1) specimens pertaining to the material originating from one type locality should form a homegeneous group (cluster); (2) material assigned to different subspecies should be represented in different clusters; (3) conspecific specimens originating from one population without their own subspecific status should join as a group one cluster formed by type specimens.

The phenograms obtained for *T. brevirostris, T. culeus, T. jelskii* and *T. rimac* do not show structures compatible with VELLARD's taxonomic suggestions. The morphometric differentiation between the five specimens of *T. brevirostris* which VELLARD assigned to three subspecies is low and the most similar pair of individuals belongs to different subspecies (R_2). The greatest morphometric difference between the 42 *T. culeus* exceeds three times that between the *T. brevirostris* specimens but none of the five morphs identified within this data set concides with any of the proposed subspecies (R_2 . 4). The same applies to the seven and five morphs, respectively, found withm the *T. jelskii* (R_2 . 5) and *T. rimac* (R_2 . 6) data sets. As objective morphometric similarly does not correspond to VELLARD's subjective grouping of conspecific speciments to subspecies, we conclude that the proposed subspecific segregation is invalid in the four species studied. This conclusion does not rule out that the definition of subspecies may be useful to describe the speciation processes due to the geographic isolation of populations of wide ranging species. However, if considered useful, the definition should be based on conceivable, objective criteria better than those presented so far.

What is the meaning of the different morphological traits (morphs = groups of conspecific specimens which are morphometrically very similar) which we detected in all seven species? As different morphs usually occur at the same locality, i.e. within the same population, they probably do not represent taxonomically relevant units. Neither do they reflect sexual dimorphism in shape, or size-specific polymorphism. The varying frequencies of morphs at different localities indicate that the adaptive value of a morph for the increase of individual fitness probably differs among the localities. Thus, intraspecific polymorphism may have evolved in response to local environmental factors such as altitude, water flow and competition. The occurrence of different morphs at the same locality may be due to temporal changes in the local environments. This interpretation is supported by the conspicious change of the frequencies of T_c -lekki morphs which took place in the Mantaro Valley during the last forty years 9 out of 10 specimens collected near Acolla in the early fifties (VELLARD, 1955), but only 1 out of 7 frogs recently collected near Acolla in the same main cluster, whereas only 1 Acolla specimen joins the remaining of Paluan specimens. Even more impressive is the same tendency in T_c rince though

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referring to geographically distant localities: only one recently collected individual pertains to the morphs dominating about forty years before (VELLARO, 1955). Attributing these changes to the different duration of preservation seems too simple because older morphs still occur (rarely) in contemporary populations, and presently dominating morphs also existed (rarely) forty years ago. Instead, there have been dramatic changes in the riparian habitats of these species during this period due to the enormous increase in human population and the resulting pollution of the rivers and streams.

In summary, we reject the proposal of subspecies for *T. brevirostris*, *T. culeus*, *T. jelskii* and *T. rimac*.

REASSESSMENT OF THE SPECIES STATUS

The large numbers of studied specimens in most species permit a reliable estimate of the ntraspecific morphometric variability and the comparison between different species. Principal component analysis of the log₁₀-transformed data did not substantially contribute to the resolution between different taxa: the usually large overlap between different species reflects their similarity in many aspects of shape. The only exception from the rule is *T bevirositis* which considerably differs from the neighbouring species (fig. 7A). However, WIENS (1993) obtained a similar low degree of resolution between eight *Telmatobias* species from northern Perú, indicating that PCA is not a powerful tool to distinguish between Andean Telmatobiinae.

In contrast, if groups (- species) are determined a priori, and if the differences between them are maximized by canonical discriminant analysis, the same morphometric data sets can distinguish between the taxa. It is noteworthy that, despite the bad image of morphometric ratios (e.g. BO0KSTEN et al., 1985), discriminant analysis using log₁₀-transformed data or ratios give very similar results and reach the same classification success. The convergent morphological adaptation to similar environmental constraints prevents an unequivocal identification at the level of individuals, but the rate of erronocusy classified individuals is low: 2. B. brachydactykas, 1.T. culeus and 1.T. jelskii out of a total of 280 specimens. WENS (1993) did not provide results of classification success in his analysis, but he also stated that discriminant analysis provided an objective base for the distinction between Andean species of Telmatobiinae. In summary, the six central Peruvian species as well as the one from southern Perú proved to be well-defined taxonomic units which posses external features allowing an objective diagnosis.

REASSESSMENT OF THE GENERIC STATUS

LAURENT (1983) recognized three genera within the central Peruvian Telmatobinas: the monotypic genera Batrachophrynus and Lynchophrys, and Telmatobius with four species. However, the evidence presented for the change of Batrachophrymus macrostomus to the genus Lynchophrys is weak: the main differences from Batrachophrymus macrostomus put forward are: (1) shorter third finger; (2) smaller size; (3) the statement that male B macrostomus lack nuptial pads. Points (1) and (2) are convergent morphological adaptations to the stream habitat which are shared with all riparian Telmatobius, whereas point (3) is simply an error (see figure 16 in Sinsecti, 1990). Moreover, both species share two morphometric features which are diagnostic for the genus Batrachophrymus despite their adaptation to different habitats: flat body and femur length exceeding tibia length Finally, the analysis of allozymes of these species and of three *Teimatobius* species (SNSCH & JURASKE, 1995) clearly demonstrates that *B. brachydactylus* and *B. macrostomus* are closely related species as originally proposed by PFIERS (1873).

In summary, we do not see any conceivable reason to split the genus Batrachophrynus and reject the proposal of Lynchophrys.

SUMMARY OF TAXONOMIC PROPOSALS

We summarized in Table VII the reassessment of the taxonomic status of central Peruvian Telmatobiinae. Three genera including seven nominal species which segregate into 15 subspecies are currently reduced to two genera *Batrachophrynus* and *Telmatobius* which include seven species without subspecific segregation.

Finally, we wish to comment on DUELLMAN'S (1993) statement that the reading of Bartachophynyuus macrostomus Peters, 1873 should be changed to B. microstomus. We do not agree because there is no doubt about the original naming by PETERS (1873), and in all research papers dealing with this species (AVLA RAMON, 1953; CAMARENA, 1953; DEDROS, 1984; FEILDAS, 1983; GORHAM, 1966; LAURERT, 1983; LAVILA, 1988B, LYNCH, 1971; 1978; MACEDO, 1950, 1960, 1976, MORALES, 1988; SINSCH, 1986, 1990; VELLARD, 1951) the original name has been used consequently.

CONVERGENT MORPHOLOGICAL TRAITS

Stream habitats require frogs to evolve morphological adaptations which enable them to move within the permanent current of water Therefore, it is not surprising that the riparian *B*. brachydactylus shares two diagnostic features with the riparian *Telmatobus*: slim head and large eye diameter. Moreover, adults of all riparan species are smaller-sized than those of the lake-inhabiting *B*. macrostomus and *T*. culves. Confusion of these convergent morphological adaptations to the same type of habitat with similar morphology due to phylogenetic relationship has led to the creation of the genus *Lynchophrys* by *LAURENT* (1983). In the Andean Telmatobinae any taxonomic conclusion based on external morphology should be backed up by other kinds of characters because the rate of erroneous classification in the morphometric distinction of the genera *Batrachophrynus* and *Telmatobius* is considerably greater than that in the distinction between stream- and lake-inhabitants: 20 % versus 8 %. Thus, the effect of convergent lines of development is probably great in this group of frogs.

GEOGRAPHICAL DISTRIBUTION

The geographical range (fig. 2) of the six central Peruvian species is still relatively unknown. We know little about the northern extension of the ranges of *T. brevirosris*, *T. carrillae* and *T. rimac*, and about the southern range of *T. jelskii*. The northern gap between the species surveyed by WENS (1993) and those in this study is subject to a recent study (SALAS, in prep.). Further attention should be paid to the exact limits of distribution of the riparian *B. brachydactylus* in relation to those of the neighbouring *Telmatobius* species. We do not even know if *B. brachydactylus* and *T. jelskii*, both present in the hydrographic system of the Mantaro river, can coexist at the same locality.

Names in use Lynchophrys brachydactyla (Peters, 1873) Batrachophrynus macrosiomus Peters, 1873		This study Batrachophrynus brachydactylus Peters, 1873 Batrachophrynus macrostomus Peters, 1873	
Telmatobius carrillae Morales, 1988 Telmatobius culeus (Garman, 1875)		Telmatobius carrillae Morales, 1988	
Termuronus cureus (Garman, 1973)	T. culeus culeus (Garman, 1875) T. culeus dispar Vellard, 1953 T. culeus secometi Angel, 1923 T culeus escul Vellard, 1951 T culeus fluviarilis Vellard, 1953 T culeus fluviarilis Vellard, 1953	Telmatobius culeus (Garman, 1875) Telmatobius culeus (Garman, 1875) Telmatobius culeus (Garman, 1875) Telmatobius culeus (Garman, 1875) Telmatobius culeus (Garman, 1875)	
Telmatobus jelskii (Peters, 1873)	T. jelsku jelski (Peters, 1873) T. jelsku bujo Vellard, 1955 T. jelski longitarsta Vellard, 1955 T jelskii walkeri (Shreve, 1941)	Telmatobius jelskii (Peters, 1873) Telmatobius jelskii (Peters, 1873) Telmatobius jelskii (Peters, 1873) Telmatobius jelskii (Peters, 1873)	
Telmatobius rimac Schmidt, 1954	T rimac rimac Schmidt, 1954 T rimac meridionalis Vellard, 1955	Telmatobius rimac Schmidt, 1954 Telmatobius rimac Schmidt, 1954	

Table VII. - Alphabetical synonymy of the telmatobune species revised in this study.

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Nevertheless, there can be little doubt that the genus *Batrachophrynus* is endemic to central Perú, as already mentioned in the original description and later in VELLARD (1951). The comment on distribution in FROST (1985) – "Andes of southern Peru and Bolivia" – is obviously wrong, as is the citation of "*Lynchophrys brachydactyla*" occurring in the northern Andes of Bolivia (De LA Riva, 1990).

CONCLUSION

At this stage of morphometric analysis we refrain from phylogenetic considerations because of the convergent adaptations of the external morphology to the same type of habitat. In the next step of analysis, we use allozyme variation within and between telmatobine species to approach the phylogenetic relationships and to compare them with phenetic relationships based on morphometry (SNESCA & JURASKE, JURASKE, JURASKE, 1995).

In conclusion, this paper provides an objective, diagnostic method to assign central Peruvian Telmatobiinae to the presently known species, based exclusively on external characters which are casy to measure. Thus, multivariate statistics, specifically canonical discriminant analysis, have proven again to be an useful tool in the classification of amphibians.

RESUMEN

Se revisa el estado taxonómico de los Telmatobina de la región central del Perú, en base a la variación intraespecífica de 18 parametros morfomètricos que presentan las sens especies (con nueve subespecies) reconocidas actualmente, las cuales se agrupan en tres géneros (Batrachophrynus, Lynchophrys y Telmatobus). Aplicando los anàlises de componentes principales y de discriminación se reconocen dos géneros (Batrachophrynus y Telmatobus) incluyendo seis especies (Batrachophrynus brachydactylus, B. macrostomus, Telmatobus) incluyendo seis especies (Batrachophrynus brachydactylus, B. macrostomus, Telmatobus brevirostris, T. carrillae, T. Jelskin, T. runac) y nunguna subespecie. Dos caracteres diagnósticos y externos distinguen las especes de Batrachophrynus y de Telmatobus, otros dos son considerados adaptaciones convergentes que diferencian a los habitantes de arroyos de aquellos que habitan las lagunas

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APPENDIX I SPECIMENS EXAMINED

Batrachophrynus brachydactylus

PERÚ: (1) Departamento Cerro de Pasco: Caza Pato, 9 males, 3 females, MHNSM 1311, 1314, 1316, 1319, 1324-1325, 1331-1336; (2) Departamento Junín: brook near Ondores: (a) first sample, 9 males, 10 females, URP 054-072; (b) second sample, 11 males, 11 females, collected by A. SALAS, March 26, 1992.

Batrachophrynus macrostomus

PERÚ: Departamento Junín: Junín Lake, Carhuamayo, 6 males, 7 females, 8 juveniles, collected by M. ANTIGNANI, February 20, 1992.

Telmatobius brevirostris

PERú: Departamento Huanuco: (1) Ambo, Chasqui, 1 male, 1 female, MHNSM 3736, 7676 (syntypes of *T. b. brevirostris*); (2) Ambo, Caina, 2 females, MHNSM 7666-7667 (syntypes of *T. b. parvulus*); (3) Santa Maria del Valle, 1 male, MHNSM 7681 (holotype of *T. b. punctatus*).

Telmatobius carrillae

PERÚ: Departamento Ancash: (1) Yuracyacu: (a) first sample, 4 males, 2 females, MHNSM 1528 (holotype), 1544-1545, 3932-3934 (paratypes); (b) second sample, 9 males,

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12 females, URP 001-021; (2) Huikia: (a) first sample, 5 males, 1 female, MHNSM 6681-6687; (b) second sample, 6 males, 2 females, URP 022-029; (3) Huaychopampa, 8 males, 4 females, URP 030-041.

Telmatobius culeus

BOLIVIA: Lake Titicaca, Isla del Sol, 1 male, 1 female, MHNSM 7769-7770 (assigned to T. c. culeus).

PERÚ: (1) Departamento Arequipa: Yura, Arequipa, 1 female, MHNSM 7678 (syntype of *t. c. exsub*; (2) Departamento Puno: (a) Azangaro, Hacienda Checayani, 1 male, 1 female, MHNSM 7673-7674 (syntypes of *T. c. lacustris*); (b) Lake Lagunillas, 6 males, 4 females, MHNSM 7768, 7776-7777, 7785-7786, 7806-7807, 7823-7825 (assigned to *t. c. escuba*; (2) Iticaca, Ocama, 3 males, 3 females, MHNSM 7779-7784 (assigned to *T. c. clacustris*); (c) Rio Have, Chucuito, 1 male, 1 female, MHNSM 7711-7772 (syntypes of *T. c. fluviatilis*); (c) Rio Have, Chucuito, 1 male, 1 female, MHNSM 7711-7772 (syntypes of *T. c. fluviatilis*); (c) Rio Have, Chucuito, 1 male, 1 female, MHNSM 7711-7772 (syntypes of *T. c. fluviatilis*); (c) Rio Have, Huuyllata, 1 male, 2 females, MHNSM 7812-7814 (assigned to *T. c. clause*), and 3 females, MHNSM 7812-7814 (assigned to *T. c. clause*), and 4 females, MHNSM 776-7767, 7787-7789, 7793-7794 (assigned to *T. c. clausris*).

Telmatobius jelskii

PERÚ: (1) Departamento Ayacucho: (a) Ayacucho, 4 males, 6 females, MHNSM 12202, 12206, 12216, 12214, 12214, 12217, 12219-12220, 12222, 12225, 12890 (assigned to T. j. walker/j; (b) Parinacochas, 6 males, 6 females, MHNSM 12838, 12841, 12883, 12901, 12904, 12910; (c) Puquio, 2 males, 1 female, MHNSM 7642-7643, 7645 (syntypes of T. j. logitarisi); (d) Tambo, 2 males, 4 females, MHNSM 7646-7651 (syntypes of T. j. lejskii bafo); (2) Departamento Huancavelica: Huancavelica, 4 males, 1 female, 2 juveniles, baf0); (2) Departamento Huancavelica: Huancavelica, 4 males, 1 female, 2 juveniles, MHNSM 7693-7641, 7660-7661, 7663-7664 (syntypes of T. j. logitarisi); (d) Departamento Junin: (a) Huancavo, Acolla, 3 males, 7 females, MHNSM 6903-6906, 6909-6914 (assigned to T. j. jelskii); (b) Huancayo, Palian, Rio Shullcas, 5 males, 2 females, 14 juveniles, collected by U. SINSCH & V. CANALES, February 19-25, 1992; (c) Tarma, Cuyrohuasi; (i) first sample, 5 males, 5 females, 1089-0075; (ii) second sample, 6 males, 5 collected by U. SINSCH & V. CANALES, February 19-25, 1992; (c) Tarma, Cuyrohuasi; (d) Fats mample, 5 males, 5 females, 1089-0075; (ii) second sample, 6 males, oflexetob y A. SALAS, March 24, 1992.

Telmatobius rimac

PERÚ: (1) Departamento Ancash: Ocros, 7 males, 3 females, MIHNSM 6935-6936, 6941-6942, 6944-6945, 6950-6951, 6953-6954 (assigned to *T. r. rimaci*); (2) Departamento Lima: (a) Canta, Obroillo, Rio Chillon, 8 males, 7 females, 7 juveniles, collected by J. ICOCHEA, March 8-9, 1992; (b) Canta, Quebrada Huaytara, 9 males, 4 females, 2 juveniles, collected by J. ICOCHEA, March 8-9, 1992; (c) Tupe, 2 males, 2 females, MHNSM 7656-7659 (syntypes of *T. r. meridionalis*).

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